

## p73 is required for endothelial cell differentiation, migration and the formation of vascular networks regulating VEGF and TGFβ signaling

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Vasculogenesis, the establishment of the vascular plexus and angiogenesis, branching of new vessels from the preexisting vasculature, involves coordinated endothelial differentiation, proliferation and migration. Disturbances in these coordinated processes may accompany diseases such as cancer. We hypothesized that the p53 family member p73, which regulates cell differentiation in several contexts, may be important in vascular development. We demonstrate that p73 deficiency perturbed vascular development in the mouse retina, decreasing vascular branching, density and stability. Furthermore, p73 deficiency could affect non endothelial cells (ECs) resulting in reduced in vivo proangiogenic milieu. Moreover, p73 functional inhibition, as well as p73 deficiency, hindered vessel sprouting, tubulogenesis and the assembly of vascular structures in mouse embryonic stem cell and induced pluripotent stem cell cultures. Therefore, p73 is necessary for EC biology and vasculogenesis and, in particular, that DNp73 regulates EC migration and tube formation capacity by regulation of expression of pro-angiogenic factors such as transforming growth factor- $\beta$  and vascular endothelial growth factors. DNp73 expression is upregulated in the tumor environment. resulting in enhanced angiogenic potential of B16-F10 melanoma cells. Our results demonstrate, by the first time, that differential p73-isoform regulation is necessary for physiological vasculogenesis and angiogenesis and DNp73 overexpression becomes a positive advantage for tumor progression due to its pro-angiogenic capacity.

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Vascular system formation is one of the earliest events during organogenesis. The original vascular plexus is established by vasculogenesis, through differentiation and assembly of mesodermal precursors.<sup>2</sup> The angiogenesis process allows the formation of new blood vessels from the existing vasculature and is perturbed in many diseases, including cancer.3 Although efforts have been made to identify factors that control vascular development, the understanding of the molecular networks remains incomplete.

The formation of new capillaries and the remodeling of preexisting blood vessels is linked by signal transduction pathways.4 The members of the p53 family (p53, p73 and p63) coordinate cell proliferation, migration and differentiation, and could act as regulators of vascular development. TP73 function in angiogenesis is quite controversial,5-7 and it has never been addressed using developmental models.

TP73 has a dual nature that resides in the existence of TA and DNp73 variants. TAp73 is capable of transactivating p53 targets<sup>8-10</sup> whereas DNp73 can act as p53 and TAp73

repressor. 11-13 TP73 final outcome will depend upon the differential expression of the TA/DNp73 isoforms in each cellular context, as they can execute synergic, as well as antagonist, functions.

TP73 role during development is emphasized by the p73knockout mice (Trp73-/-, p73KO from now on) multiple growth defects.<sup>14</sup> These mice, which lack all p73 isoforms, exhibit gastrointestinal and cranial hemorrhages, 14 suggestive of vascular fragility. Furthermore, TAp73 directly regulates GATA-1,8 which is essential for endothelial and hematopoietic differentiation. 15,16 This compounded information led us to hypothesize that p73 could be implicated in the regulation of vasculogenesis/angiogenesis.

Regulation of these processes involves a broad range of signaling molecules essential for vascular growth and stability, 17 such as vascular endothelial growth factor (VEGF)<sup>18</sup> and transforming growth factor- $\beta$  (TGF- $\beta$ ).<sup>19</sup> TGF- $\beta$ operates as a rheostat that controls endothelial cell (EC) differentiation, having an inhibitory effect on EC migration and proliferation by the TGF-β/TGFRI (ALK5)/Smad2/3 pathway,

Abbreviations: EB, embryoid body; EC, endothelial cells; HUVEC, human umbilical vein cells; iPSC, induced pluripotent stem cells; mESC, mouse embryonic stem cells; p73KO, p73 – / – ; TGF- $\beta$ , transforming growth factor- $\beta$ ; VEGF, vascular endothelial growth factor; WT, wild type Received 15.4.14; revised 12.11.14; accepted 13.11.14; Edited by E Candi; published online 09.1.15

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while the TβRII-ALK5/ALK1 complex activates Smad1/5/8, ID1 expression and a pro-angiogenic state. 20-22

Regulation of the TGF- $\beta$  and VEGF pathways by p53 family members has been documented.<sup>23,24</sup> However, p73's function in these pathways during development remains largely unexplored. In this work, we have used mouse embryonic stem cells (mESC) and induced pluripotent stem cells (iPSCs) as models that recapitulate early vascular morphogenesis.<sup>25–27</sup> ESC and iPSC form multi-cellular aggregates (embryoid bodies, EBs) that, under appropriate conditions, generate functional EC.28 mESC and iPSC differentiation capacity into ECs has addressed. 29,30 We have also performed retinal vascularization analysis to assess vascular processes in vivo. 31,32

We demonstrate that p73 deficiency perturbs density and stability of mouse retinal development by affecting VEGF and TGF- $\beta$  signaling. Furthermore, p73 is necessary for the assembly of vascular structures under physiological conditions in mESC and iPSC. Moreover, DNp73 positively affects angiogenesis through regulation of the TGF-β pathway in human umbilical vein cells (HUVEC) and DNp73overexpression results in enhanced angiogenic potential of B16-F10 melanoma cells.

## Results

Defects in retinal vessel migration and vascular morphology in p73-deficient mice. P5-p73KO and wildtype (WT) mice retinas were analyzed to ascertain p73 developmental function in vascular formation using isolectin B4 (IB4) staining. At the leading edge of the vascular plexus (sprouting zone), WT-tip cells and their extending filopodia were directed toward the periphery, guiding the direction of vascular growth (Figure 1a, arrows). In contrast, p73KO retinas had disoriented tip cells (arrows) with long and thin filopodia (Figures 1a, circles and 1b). The central plexus was also abnormal, appearing disorganized, less dense and branched than WT (Figure 1a, right panel). Morphometric analysis showed increased lacunarity and moderate, but highly significant, decrease in vessel covered area, vessel length and branching index, indicating a less dense p73deficient vascular plexus (Figure 1b).

P5 and P7-p73KO vascular plexus had less spreading suggesting that the abnormal filopodia was affecting tip cell migration<sup>33</sup> (Figure 1c). IB4 positive (IB4+) macrophages, which act as cellular chaperones for EC fusion, 34 appeared as

isolated cell clusters, rather than interacting with tip cell filopodia (Figure 1a, middle panel, arrow heads), supporting the idea of a defective guidance and migration in p73deficient ECs.

We sought to analyze whether p73-loss affected vessel architecture and stability. The vascular basement membrane, marked by type IV-collagen staining, is closely associated with the EC abluminal surface. 35 Thus, empty basement membrane sleeves (collagen-IV+/IB4-) identify vessel regression.<sup>36</sup> In WT central plexus, most vessels showed collagen-IV+/IB4+ staining with only a few empty collagen-IV sleeves (Figure 1d, arrows). However, in p73KO, empty sleeves were frequent, suggesting that p73 deficiency results in vessel regression.

Next, we examined the resident astrocytic network, which provides guidance cues for ECs by establishing a VEGF-A gradient. Although in WT retinas, astrocytes and ECs closely followed one another, in p73-deficient retinas astrocytes displayed a chaotic reticulation with astrocytes clumps (Figure 1e, white arrows) surrounded by IB4+cells (arrow heads). p73KO retinas had significantly reduced VEGF-A expression and distribution (Figures 1f, yellow arrows and 2a), confirming a hindered pro-angiogenic signaling in the absence of p73. WT retinas showed IB4+/VEGF-A+-microglial cells commonly found where two tip-cells were contacting each other, at sites of prospective sprout anastomosis (Figure 1f. arrow heads). In p73KO, microglia was isolated and not fully interacting with the ECs. Thus, lack of p73 affects the retinal microglia, hindering the VEGF-A gradient for EC migration.

 $TGF-\beta$  signaling is essential for EC organized migration and determines EC function during development. 20,37 We detected a significant decrease in TGF-β1, TGF-βR1 and ALK1 levels in p73KO retinas (Figures 2b-d), and significantly lower ID1 levels (Figure 2e), suggesting an attenuated pro-angiogenic state. Taken together, our data demonstrate that p73 is necessary for retinal vasculature organization and suggest that p73 could be a positive regulator of EC migration, vessel sprouting and tubulogenesis in vivo regulating the VEGF and TGF- $\beta$  pathways.

p73 deficiency results in impaired endothelial differentiation and reduced angiogenic sprouting in mESCs. We used mESC and iPSC to address p73 function in EC differentiation and vasculogenesis. Both TA and DNp73 isoforms were expressed in mESCs at low but consistent levels, with DNp73 being the most abundant isoform

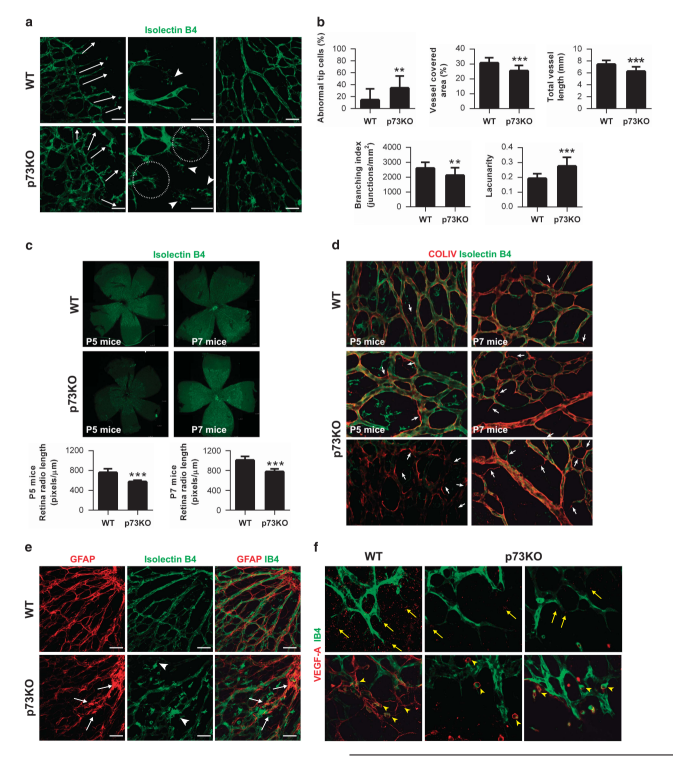
Figure 1 Absence of p73 perturbs development of the retinal vasculature. (a) Retinas from P5 WT and p73KO mice were stained with isolectin B4 (IB4) to analyze the morphology and orientation (white arrows, left panels) of filopodia at the leading edge of vascularization. Higher-magnification images for each genotype (medium panels) show abundant and disorientated filopodia (circles) and tissue macrophages (arrow heads) in p73KO retinas. Scale bars = 50 \( \textit{\pm} m. \) Right panels illustrate perturbed vascular branching and decreased vascular density in p73KO central retinal plexus compared with WT. (b) Quantification of vessel coverage (percentage of area covered by IB4\* endothelial cells), total vessel length, vascular branching index (branch points/unit area) and lacunarity (distribution of the gap sizes surrounding the object). Representative images were analyzed independently using the AngioTool software (https://ccrod.cancer.gov/confluence/display/ROB2/Home). (c) IB4 retinal flat-mount staining of P5 and P7 retinas. Radio was measured from the optic nerve to the sprouting zone. The spreading of the vasculature toward the periphery is highly significantly reduced in p73KO retinas. (d) Visualization of empty matrix sleeves by IB4 (green) and collagen IV (red) staining, with increased presence of collagen IV sleeve segments lacking endothelial cells (IB4-negative; white arrows) in p73KO retinas. (e) GFAP/IB4 double staining to visualize astrocytes (red) and vasculature (green), respectively. p73KO retinas display a disorganized astrocyte network underlying a chaotic vasculature; tufts are indicated by arrow heads. (f) VEGF-A immunostaining (red) in P5 retinas. Note that VEGF-A expression is markedly decreased in the absence of p73. Yellow arrow heads indicate IB4+/VEGF-A+ microglial cells commonly found at sites of prospective sprout anastomosis. All statistical analysis were performed with data from at least five animals. Bar graphs represent mean ± S.D. Equal-variance Student's t-test was performed to evaluate statistical differences. \*\*P < 0.01; \*\*\*P < 0.001



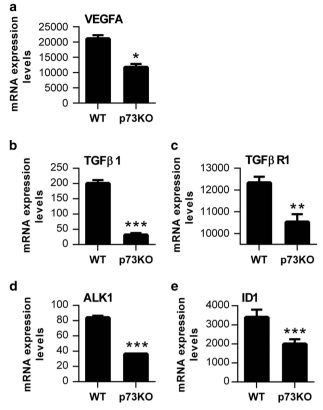
(Figure 3a). Stable cell lines with constitutive p73-dominant-negative mutant (DDp73) expression were generated<sup>38</sup> (Supplementary Figure 1a). The clones exhibited the typical ES morphology, alkaline phosphatase activity as well as Nanog expression (Supplementary Figure 1b), and were able to differentiate into derivatives of the three germ layers (Supplementary Figure 1c). Although DDp73 cells expressed the mesodermal/endothelial marker CD31, they failed to form vascular networks like the WT cells (Supplementary Figures

1c and d), supporting a possible role for p73 in endothelial morphogenesis.

Next, we differentiated mESCs under two-dimensional (2D) and three-dimensional (3D) endothelial-specific conditions (Supplementary Figure 1e). The 3D model recapitulates the developmental steps of vasculogenesis and sprouting angiogenesis, whereas the 2D EBs recapitulate the formation of the primitive vascular network *in vivo*.<sup>25</sup> EBs were immunostained for CD31 and vascular endothelial (VE)-cadherin and EB size







**Figure 2** Lack of p73 affects TGF-β signaling *in vivo*. Quantification of VEGF-A expression and analysis of the TGF-β/ALK1/ID1 signaling pathway in P5 retinas from WT and p73KO mice. qRT-PCR analysis demonstrated a significant decrease in VEGF-A expression (**a**), TGF-β1 expression (**b**), TGF-βR1 expression (**c**), ALK1 expression (**d**) and ID1 expression levels (**e**) in the absence of p73. Analysis was performed with data from three independent experiments. Mean  $\pm$  S.D. are represented; equal-variance Student's *t*-test was performed to evaluate statistical differences. \*P<0.05, \*\*P<0.01, \*\*\*P<0.001

and morphology were analyzed (Figures 3b and c). Control EBs had CD31<sup>+</sup> cells forming a branching network from the center of the EB to the outer rim (Figure 3b, upper panel), whereas DDp73-EBs had a smaller diameter and their ECs (CD31<sup>+</sup> or VE-cadherin<sup>+</sup>) did not assemble into a vascular network (Figures 3b and c).

In 3D conditions, control EBs formed a vascular network after 7 days in *vitro* (7 DIV) whereas DDp73-EBs failed to sprout (Figure 3d). After 10 DIV, more than 75% of control EBs showed branched CD31<sup>+</sup> and VE-cadherin<sup>+</sup> sprouts (Figures 3e and f). However, DDp73-EBs had fewer and shorter sprouts that failed to form a branched network. In DDp73-EBs only some VE-cadherin<sup>-</sup> cells migrated away from the EB core (Figure 3f, arrows), supporting the notion that p73 deficiency affects EC migration and tube formation.

DDp73 effect on vascular morphogenesis may involve a crossed effect on other p53 family members. <sup>38</sup> To rule out this possibility, iPSC reprogrammed from p73KO or WT mouse embryonic fibroblasts were used (Martin-Lopez *et al.*, manuscript under preparation). WT-iPSC-EBs in 3D culture exhibited abundant sprouts that constituted an intricate network (Figures 3g and h). In contrast, only few p73KO-iPSC-EBs had sprouts that were short and unable to form networks (Figure 3g). CD31-immunostaining highlighted the lack of

sprouts of p73KO-iPSC-EBs (Figure 3h). To rule out the possibility that reprogramming had affected the genetically modified iPSCs capacity to form 3D vascular sprouts, we also differentiated p53KO and p73KO/p53KO iPSC. p53KO-iPSC EBs generated a profuse vascular network, demonstrating that reprogramming did not affect the capacity of KO cells to differentiate into ECs and form vascular sprouts. Thus, p53 was not required for this process, but lack of p73, even in the context of p53-deficiency, resulted in lack of vascular sprouts (Supplementary Figure 2a), confirming p73 requirement and that the defect in sprout formation was not due to enhanced p53-dependent cell death.

To address whether p73 deficiency could have a non-cell autonomous effect, we performed a co-culture experiment mixing GFP+-mESC with either WT-iPSC or p73KO-iPSC, and differentiating them under 3D endothelial-specific conditions (Supplementary Figure 2b). Although GFP+-mESC/WT-iPS EBs were able to differentiate into ECs that form vascular sprouts, only few GFP+-cells from the GFP+-mESC/p73KO-iPS EBs were capable to migrate and form sprouts. These results supported p73 critical role in vascular morphogenesis, but also indicated that p73 deficiency has a non-cell autonomous effect over neighboring WT-ECs.

To study the mechanism of p73 regulation on EC commitment, endothelial CD31+ cells and differentiated (but nonendothelial) CD31<sup>-</sup> cells from 12 DIV-EBs were isolated. The number of CD31+ cells from DDp73-EBs was significantly lower than from control-EBs (Figure 4a), indicating that p73 function is required for efficient endothelial differentiation. To determine the contribution of p73 isoforms in this process, we compared TA and DNp73 expression in undifferentiated mESC (gray bars) with that in differentiated CD31+ ECs (black bars). Both isoforms were significantly upregulated in the CD31<sup>+</sup> population compared with the undifferentiated mESC, with DNp73 being predominant in ECs (Figure 4b). The nonendothelial differentiated CD31- cells (white bars) showed the opposite pattern of isoform expression (TA > DN). Analysis of EC markers revealed that CD31 and VE-cadherin levels were significantly lower in DDp73-CD31+ cells (Figure 4c), confirming that p73 functional inhibition impairs EC differentiation.

So far our results suggest that p73 could be regulating some of the molecular pathways that control endothelial differentiation, migration and/or assembly of endothelial precursors. To address this, we analyzed VEGF-A and TGF- $\beta$  signaling, both altered in p73KO retinas. In agreement with previous publications, <sup>6,7</sup> DDp73-CD31<sup>+</sup> cells expressed lower levels of VEGF-A (Figure 4d). The expression of VEGFR1, a decoy of angiogenic signaling, was significantly higher in DDp73-CD31+, while VEGFR2 expression, the VEGF signaling transducer, was lower (Figure 4d). The VEGFR2/VEGFR1 ratio, which may be considered as a pro-angiogenic indicator, 39 was about five times lower in DDp73-CD31+. Furthermore, we detected a significant reduction in TGF-B expression, with a concomitant attenuation in ID1 levels, in DDp73-CD31<sup>+</sup> cells (Figure 4e), all consistent with a decrease in the number of cells committed to endothelial lineage. 40

Taken together, our data indicate that the lack of p73 in mESC attenuates endothelial differentiation, diminishes the angiogenic potential of the differentiated cells and hinders angiogenic sprouting of 3D-EBs, therefore demonstrating that

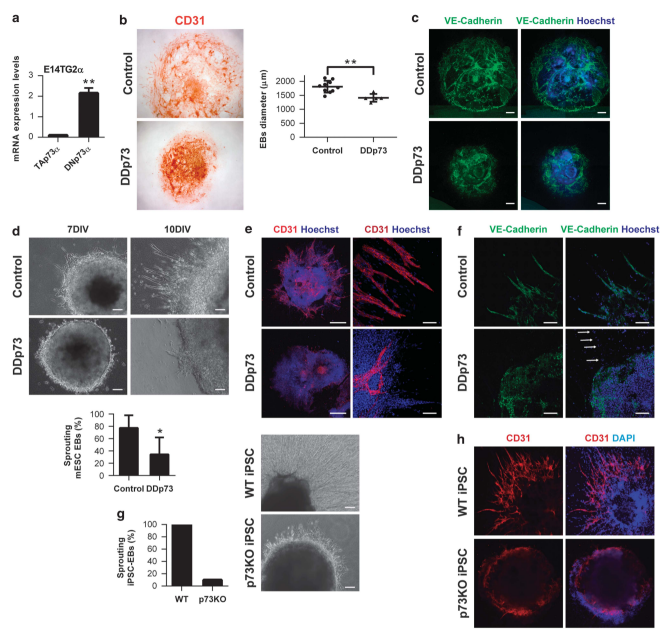


Figure 3 Lack of p73 impairs the formation of vascular structures and endothelial sprouting of mESC and iPSC in the EB endothelial differentiation model. (a) Quantitative analysis of p73 isoform expression (TA and DN) by qRT-PCR in proliferating E14TG2a mESC. Ct-TAp73: 38.22 and ΔCt-TAp73: 0.099; Ct-DNp73: 33.78 and ΔCt-DNp73: 2.15 (b−c) EBs size, morphology and formation of vascular structures in WT and DDp73 mESC EBs under 2D differentiation culture conditions. (b) EBs were stained for CD31 expression and EB average diameter for individual EBs ( $n \ge 10$ ) was calculated after 7 days *in vitro* (7 DIV). (c) VE-cadherin immunostaining (green) shows that DDp73-EBs did not assemble into vascular networks. (d−f) Angiogenic sprouting in WT and DDp73 mESC EBs under 3D differentiation conditions. (d) Phase contrast images correspond to 7 DIV and 1D DIV EBs. Graph shows percentage of sprouting EBs at day 18. (e) CD31 expression (red) and (f) VE-cadherin expression (green) demonstrate that DDp73 mESC EBs inder 3D differentiation conditions. (g) Phase contrast images of 12 DIV iPSC-EBs illustrate sprouts in WT EBs, but not in the p73KO-iPSC-EBs. Graph represents percentage of sprouting EBs at day 18. (h) CD31 expression (red) highlighted that p73-deficient iPSC cannot form vascular structures. Data represent mean values ± S.D.; equal-variance Student's Ftest was performed to evaluate statistical differences. \*P<0.05, \*\*P<0.01

p73 is a positive regulator of *in vitro* vasculogenesis/ angiogenesis, at least in part, by modulating pro-angiogenic signaling through the VEGF and TGF- $\beta$  pathways.

**DNp73** silencing interferes with HUVECs to undergo migration and tubulogenesis. To show that p73 has a specific role in EC biology, independently of its possible effect

in other cell types within the retina or in the EBs, we used HUVECs. p73 isoforms were differentially expressed in HUVECs, with DNp73 expression being about 2.7 times higher than TAp73 (2.7  $\pm$  0.0002; Ct-TAp73: 36.70 and  $\Delta$ Ct-TAp73: 0.00019; Ct-DNp73: 35.60 and  $\Delta$ Ct-DNp73: 0.00045). We suppressed p73 function with DDp73 or by an RNA interference knockdown (p73KD) using previously validated

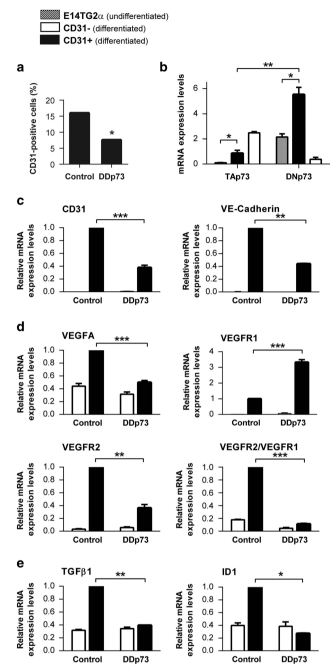


Figure 4 p73 functional inhibition hinders mESC endothelial differentiation, decreases endothelial marker expression and blunts the pro-angiogenic state of the obtained endothelial cells. (a) CD31 positive cells were isolated with magnetic beads and the obtained cell number was quantified. (b) qRT-PCR analysis of p73 isoform expression in proliferating undifferentiated E14Tg2a (square bars), differentiated nonendothelial CD31- negative cells (CD31-, white bars) and differentiated CD31 positive endothelial cells (CD31+, black bars) demonstrated that TA and DNp73 are differentially regulated during mESC endothelial differentiation. (c-e) Quantitative gRT-PCR analysis of endothelial cell marker expression (c), VEGF signaling (d) and TGF-β/ID1 signaling (e), in CD31<sup>-</sup> (white bars) and CD31<sup>+</sup> cells (black bars) isolated from either control or p73DD-EBs. Bars represent mean values ± S.D.; experiments were repeated twice; equal-variance Student's t-test was performed to evaluate statistical differences. \*P < 0.05, \*\*P < 0.01, \*\*\*P < 0.001

small interfering RNA (siRNA) oligos: p73i.4 for total-p73KD, TAp73i for specific TAp73 or DNp73i for DNp73-KD.8 The maximal interference using these siRNAs occurs 72 h after transfection, lasting more than 48 h.8 All angiogenic assays were carried out during this period. It is important to point out that p73KD did not affect HUVEC proliferation kinetics within this time frame (Supplementary figure 3).

HUVECs were transfected with either DDp73 or siRNAip73i.4 and tube formation was monitored and quantitatively analyzed, p73-deficient HUVEC showed reduced tube formation capacity and networks with reduced area and fewer branching points and loops (Figures 5a and b). Total-p73KD and DNp73 in particular, but not TAp73 alone, provoked a significant reduction in all the analyzed parameters (Figure 5c). These parameters revealed a defective capacity of the interfered ECs to migrate and assemble into tubular structures, indicating that DNp73 deficiency results in an impaired endothelial morphogenesis. On the contrary, under normoxic conditions, there was no significant effect on any of the analyzed parameters after TAp73 KD.

Next, p73 role in EC migration was addressed by wound healing or 'scratch' assays.41 The percentage of wound closure was significantly reduced upon functional inhibition or total p73KD (Figure 6a). DNp73KD, but not TAp73KD alone, attenuated wound closure (Figure 6c), indicating that DNp73 is a positive regulator of EC migration. We examined TA and DN expression kinetics after the wounding. As shown in Figure 6b, both isoforms were upregulated to a similar extent, indicating that the EC-characteristic DNp73>TAp73 ratio was maintained. To conclusively demonstrate that DNp73 function was necessary for EC migration, we performed a phenotype recovery experiment. DNp73 overexpression in siRNA-p73i4 cells restored the migratory capacity (Figure 6d), confirming DNp73 as a positive regulator of EC migration.

To elucidate the molecular mechanisms underlying the DNp73 regulation, we analyzed the VEGF and TGF-β pathways. We examined phosphorylated ERK1/2 as a downstream indicator of VEGF signaling<sup>42</sup> and, as published,<sup>43</sup> both p73 isoforms regulated this pathway (Figure 7a). VEGF-A levels were strongly downregulated upon total p73 and DNp73 silencing, (Figures 7a and b), and barely affected after TAp73KD (Figure 7b), pointing to DNp73 as the predominant regulator on HUVEC normoxic cultures. However, transcriptional analysis indicated that ectopic DNp73 expression did not induce VEGF-A promoter activity (Supplementary Figure 4a), suggesting the existence of other downstream posttranscriptional regulatory mechanisms.

On the other hand, analysis of TGF-β1/ALK1/ID1 pathways showed that only DNp73 is upregulated by TGF-β1 treatment in a dose-dependent manner (Figure 7c), linking p73 differential regulation to the angiogenic TGF-β1 signaling axis. Indeed, DNp73 silencing resulted in significantly reduced p-SMAD1/5 and total SMAD1 proteins (Figure 7a), and TGFβ1, ALK1 and ID1 levels (Figure 7d), unequivocally demonstrating that DNp73 is required for TGF-β1/ALK1/ID1 signaling in ECs.

We have demonstrated p73 requirement for EC biology and vasculogenesis and, in particular, that DNp73 regulates EC migration and tube formation. However, our in vivo data also indicated that p73 deficiency could affect non-EC resulting in

reduced pro-angiogenic milieu. We sought to address whether either silenced (Supplementary figure 4b) or transiently DNp73 could affect tumor angiogenesis, using the syngeneic transplantation model of B16-F10 melanoma.<sup>44</sup> Conditioned media (CM) was prepared from B16-F10 melanoma cells with overexpressed DNp73 (Supplementary figure 4c), and its angiogenic effect on HUVEC was tested in scratch assays. DNp73-deficient CM retard, but do not suppress, HUVEC